PATENT SPECIFICATION

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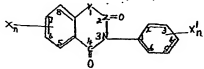
COMPLETE SPECIFICATION

Benzoazinediones and Germicidal Compositions made therewith

We, STECKER INTERNATIONAL S.P.A., a body corporate organised under the laws of Italy, of Via Turati No. 29, Milan, Italy, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:

This invention relates to the preparation of new benzoazinediones, including benzo-

This invention relates to the preparation of new benzoazinediones, including benzothioxazinediones and benzoxazinediones, and to novel germicidal compositions prepared therewith. The compounds which are the subject of the present invention fall within the generic formula:



where X and X1 are chlorine, bromine, iodine or CF₁,

n is an integer from 0 to 3, subject to the proviso that X or X¹ represent at least one and not more than two CF, groups,

Y is sulphur or oxygen, and

Z is sulphur or carbon.

The small numerals within the nuclei are inserted merely for more convenient orientation of the derivatives to be discussed herein.

The compounds of the present invention may be prepared by reacting a substituted salicylanilide with thionyl chloride, phosgene, or ethyl chloroformate according to the following typical reactions:

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In reaction (1) 3,5-dibromo-3-(trifluoromethyl) salicylanilide is reacted with thionyl chloride to produce 6,8-dibromo-3-(3-trifluoromethylphenyl)-1,3-benzothiox-azine-2,4-dione. In reaction (2) the same salicylanilide is reacted with ethyl chloroformate to produce 6,8-dibromo-3-(3-trifluoromethylphenyl)-1,3-benzoxizine-2,4-dione.

These compounds may be prepared according to the method described by Stanseth, Baker and Roman, J. Med. Chem., 6, 1212 (1963). A typical method of preparation is as follows:

6,8-Dibromo-3-(3-trifluoromethylphenyl)-1,3-benzoxaxino-2,4-dione.

A molal solution of 3,5-dibromo-3'-(trifluoromethyl) salicylanilide in a mixture of A motal southon of 3,3-dioromo-3 (trinuoromethyl) sancytaninde in a mixture of pyridine and acetonitrile is stirred at 2-5°C, during dropwise addition of a motal quantity of ethyl chloroformate. Stirring is continued for 1-2 hours while the temperature is gradually increased to 120°-125°C. After about 60 mls. of distillate has been collected in a Barrett trap, the mixture is slowly cooled and, before it is solidified, water and concentrated HCl are added with stirring and further cooling. The crude product is then isolated, washed with water, and air-dried. The compound may be recrystallized from acetone, after decolorization with activated charcoal. The recrystallized product is then recovered.

Table I gives a list of compounds which have been prepared in accordance with

the foregoing method.

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Š	Salicylanilide ·	Resciant	Product	Properties
	3,5-Dibromo-3'-(trifluoromethyl)	SO CI,	6,8-dibromo-3-(3-trifluoromethyl- phenyl)-1,3-benzothioxazine-2,4-dione	ш.р. 190—5°С.
	3,5-Dibromo-3'-(trifluoromethyl)	Bthyl chloroformate	6,8-dibromo-3-(3-trifluocomethy)- phenyl)-1,3-benzozazine-2,4-dione	mp. 233—5°C
_	3'-(trifluoromethyl)	Ethyl chloroformate	3-(3-trifluoromethylphenyl)-1,3- benzoxazine-2,4-dione	пр. 198—199°С
	2'-Chloro-3'-(triffuoromethyl)	Ethyl chloroformate	3-(2-chloro-3-trifinoromethylphenyl)- 1,3-benzoxazine-2,4-dione	щр. 195—198°С.
	3,5-Diodo-3',5'-bis(trifluoromethyl)	Ethyl chloroformste	6,8-diiodo-3-(3,5-bis(trifluoromethyl- phenyl)-1,3-benzozzazne-2,4-dione	m.p. 214—8°C.
	2-Thiopbenyl-3,5-dibromo-3'- (riffuoromethyl)	Ethyl chloroformate	6,8-dibromo-3-(3-trifluoromethylphenyl)-1,3-benzothiazine-2,4-dione	m.p. 238—40°C.
	3,5-Dichloro-4-(trifluoromethyl)- 4'-iodo	Ethyl chloroformate	6,8-dichloro-7-(trifluoromethyl)-3 (4-lodophenyl)-1,3-benzozazine- 2,4-dione	H.p. 220—4°C

	The compounds of the present invention have been found to show unexpectedly	
	high toxicity to micro-organisms, such as bacteria, rough, and organisms. The antibacterial	
	compared to the unsubstituted of heartful to the unsubstituted of heartful to the present compounds is shown in Table II, as the minimum inhibitory activity of the present compounds is shown in Table II, as the minimum inhibitory activity of the present compounds is a 24-boar booth culture of each	5
5	The second of th	,
	organism was made in Brain Heart Intustor, making organism 9 mls. of BHI broth, were	
	A number of screw cap test times, each containing of 100 ml. prepared and sterilized for 15 minutes at 25 psi at 120°C. A number of 100 ml. prepared and sterilized for 15 minutes at 25 psi at 120°C. A number of 100 ml.	
	prepared and sterilized for 15 indicates at 25 per BHI broth also were prepared, were volumetric flasks, each commaning about 80 ml. of BHI broth also were prepared, were volumetric flasks, each commaning about 30 ml. of BHI broth also were prepared, were	10
10	capped with glass caps, and were stermeet in the tested was accurately weighed and	
	One tenth of a gram of the compound to be tested was transferred with aseptic was dissolved in acctone or alcohol. The mixture then was transferred with aseptic was dissolved in acctone or acceptance of the compound to be tested was transferred with aseptic was transferred with aseptic was transferred with a septic was transferred wit	
	was dissolved in acctone or alcohol. The mixture that was containing the BHI broth, technique to the previously sterilized volumetric flasks containing the BHI broth, technique to the previously sterilized volumetric flasks containing the BHI broth. This	
	With aseptic technique, the mixture was not the compound to be tested.	15
15	With aseptic technique, the inixture was attended to be tested. mixture then consisted of 1:1000 dilution of the compound to be tested. Ten ml. of this mixture were transfered aseptically to a sterile capped tube by	
	Ten ml. of this mixture were transfered ascendary to a state of the stock solution with a 10 ml. Mohr pipette. Serial dilutions then were made from this stock solution with a 10 ml. Mohr pipette. Serial dilutions then were made from this stock solution with	
	concentrations of 1:10,000, 1:100,000, 1.1,000,000 and 110,000,000	
	pounds	20
20	To each of the dilutions of a given compount that relatively of the broth solutions hour broth culture of the organism to be tested. The unfoldity of the broth solutions hour broth culture of the organism to be tested. The density of the broth solutions	
	was determined by a Weish Densiciant The broth	•
	observation for purposes of accuracy with the at 270°C A control consisting of 0.1 ml.	25
25	dilutions were then allowed to stand for 24 lift. at 15 orth also was prepared and was of a 24-hour broth culture and 9 ml. of BHI broth also was prepared and was of a 24-hour broth culture and 9 ml. of BHI broth also was prepared and was	25
	of a 24-hour broth culture and 9 int. of 1811 both bettered. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected by the compounds to be tested. At the end of the subjected by the compounds to be tested. At the end of the subjected to the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested. At the end of the subjected to the same conditions as the compounds to be tested.	
	24-hour period, the tubes again were observed in turbidity in the broth. occurred, it would be manifested by an increase in turbidity in the broth.	
	occurred, it would be manifested by an increase in turbidity in the state antibacterial activities All compounds were subsected to the same testing and their antibacterial activities	30
30	were compared.	30
	The unsubstituted compound No. 1 of Table by the present invention. The expression ineffectiveness, as compared to the compounds of the present invention. The expression ineffectiveness, as compared to the compounds inhibiting and killing action against	
	ineffectiveness, as compared to the compounds of the present invention against "germicidal or antibacterial activity" includes inhibiting and killing action against germicidal or antibacterial activity" includes inhibiting and killing action against	
	bactera, fungi and similar organisms. The const R coli L. casei, and others.	35
35	been found effective against organisms such as 3. syphs, or more com- The present germicides are useful in compositions comprising one or more com-	
	The present germicides are useful in compositions commission, i.e., relatively pounds of the present invention and a germicidally inert material, i.e., relatively pounds of the present inventional diluent same and/or detergent, and plastics	
	speaking, such as an inert international adventure of the impregnated with one	
40	and/or rubber. Fibrous materials may also advantageously, some soaps and detergents or more compounds of the present invention. For example, some soaps and detergents or more compounds of the present inventions relative to those of the compounds of	40
40	possess a bactericidal action, but such a titled, effect in comparison with the overall	
	the present invention, is weak and of the compositions the compounds of the	
	germicidal activity of the composition. In such compositions as low as 10 p.p.m., although, present invention may be employed in concentrations as low as 10 p.p.m. or 0.001%	AE
45	present invention may be employed in concentrations as so p.p.m. or 0.001% from a practical point of view, it is desirable to use as much as 50 p.p.m. or 0.001% from a practical point of view, it is desirable to use as much as 100 or 5%, or even more.	45
	by weight, or 0.01%, 0.1%, 0.5% of as interest importion are those comprising	•
	Particularly useful compositions of the present inventor the soaps and detergents, and especially toilet soaps of cosmetic detergents in which the soaps and detergents, and especially toilet soaps of cosmetic detergents in which the	
	compounds of the present inventor may be a "determent" employed	50
50	0.1%, 0.5% or even up to 11% by weight, a second elementary compositions including	
	herein will be used to include all synthetic and natural ceaning constraints are mide-propyl-2-hydroxy-ammonium cationic detergents, such as dimethyl stearamide-propyl-2-hydroxy-ammonium cationic detergents, such as dimethyl stearamide-propyl-2-hydroxy-ammonium	
	dihydrogen phosphate, amone detagents, san a fetty and similar acids, e.g.,	
	soaps of hydrolyzed natural or syllakate by the determine such as sarcosine.	55
55	sodium and porassium stearates or oleates, ampnoyite detergents, natural non-ionic detergents, such as polyoxypropylene polyoxypethylene condensates, natural non-ionic detergents, such as polyoxypropylene polyoxypethylene condensates, natural non-ionic detergents, such as polyoxypropylene pums, and the mixtures thereof. The term	
-	detergents, sign as starting, regulating general meaning i.e. a cleansing	
	"sosp" employed ferent is used in a popular compound, such as potassium or sodium	60
60	hydroxide and a fat or fatty acid, both saturated and unsaturated, hydroxide and a fat or fatty acid, both saturated and unsaturated.	60
Ų.	Another valuable use of the compounds of the provides and paper oulds	
	to sanitize fibrous material, such as continuous A so, by weight They also serve as	
	preferably in concentrations of about 0.01% to 0.5% by weight prior to antiseptic agents, when incorporated in plastic or rubber compositions, prior to	

molding into articles of commerce, such as baby rattles, gloves, and food wrappers, preferably in concentrations of 0.0051% to 0.51% by weight.

TARE II

	Compound	Effectiveness Against S. aureus MIC × 10*
1	3-Phenylbenzonazine-2,4-dione	1:1 — 1:10
	6,8-Dibromo-3(3-trifinoromethylphenyl)-1,3-benzaszine-2,4-dione	1:1000 — 1:10,000
	3-(3-Triffuoromethylphenyl)-1,3-benzoxazine-2,4-dione	1:100 — 1:1000
	3-(3-Trifluoromethyl-2-chloro-phenyl)-1,3-benzozazine-2,4-dione	1:1000 — 1:10,000

WHAT WE CLAIM IS:—
1. Compounds having the general formula:

$$X_n - X_n = 0$$
 $X_n = 0$
 $X_n = 0$

5	where: X and X ¹ are chlorine, bromine, iodine, or CF ₂ , x and X ¹ are chlorine, bromine, iodine, or CF ₂ , n is an integer from 0 to 3, subject to the proviso that X or X ¹ represent at least one and not more than two CF ₂ groups,	5
1Õ	Y is sulphur or oxygen, and Z is sulphur or carbon. Compounds according to claim 1, wherein Y is oxygen. Compounds according to claim 1, wherein Y is oxygen and Z is carbon, and	10
15	wherein n is an integer from 1 to 3. 4. Compounds according to claim 1 wherein Z is sulphur. 5. Compounds according to claim 1 wherein Y is sulphur and wherein n is an integer from 1 to 3.	IJ
D	6. Compounds according to claim 1 wherein 1 is oxygen, 2 is oxygen, 2 is wherein n is an integer from 1 to 3.	
20	8. The compound 6,8-dibromo-3-(3-trimoromethylphenyl)-1,3-dichloro-4-trifluoromethylphenyl)-1,3-	20
25	benzothiazine-2,4-dione. 10. Compositions comprising at least one compound according to any of the preceding claims, together with an inert pharmaceutical diluent. 11. Compositions comprising at least one compound according to any of claims 1 to 9 together with a soap and/or detergent, both as hereinbefore defined. 12. Compositions comprising at least one compound according to any of claims	25
	1 to 9 together with plastics and/or rubber. 13. Fibrous materials whenever impregnated with at least one compound accord-	30
30	ing to any of claims 1 to 9. 14. Compositions according to claim 11 wherein the total weight of said compounds is in the range 0.001% to 5% of the total weight of the composition. 15. Compositions according to claim 12 wherein the total weight of said compounds is in the range 0.005% to 0.5% of the total weight of the composition.	35
35	16. Fibrous materials according to claim 15 whether to the compounds is in the range 0.01% to 0.5% of the total weight of said impregnated	33
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-	5, Park Gardens, Glasgow, C.3.	

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